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SOLUBILIZATION METHOD OF DYE INTO AROMATIC HYDROCARBON SOLVENT [Senryou no houkouzoku tankasuiso kei youzai eno kayouka houhou]

Isao Niimura, et al.

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INVENTOR(S)	(72):	NIIMURA, ISAO; MAEDA, SHIGEO; HOSHINO, KENICHI; SAKAI, TAKAYUKI
APPLICANT(S)	(71):	HODOGAYA KAGAKU KOGYO CO.
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1. Title of the Invention

SOLUBILILZATION METHOD OF DYE INTO AROMATIC HYDROCARBON SOLVENT

2. Claims

A solubilization method of a dye into an aromatic hydrocarbon solvent characterized by a mixing treatment of said dye with a polymer prepared from polypropylene glycol and ethylene oxide having a general formula as shown below:

(wherein in the above formula, n is integers of 15 \sim 50; m + m' is integers of 5 \sim 30; and the sum of m + m' + n is 20 \sim 80).

3. Detailed explanation of the invention

The present invention relates to a solubilization method of a dye into an aromatic hydrocarbon solvent such as benzene, toluene, xylene, cymene, naphthalene, styrene and the like.

Explaining in more detail, the present invention relates to a solubilization method of a metal complex salt dye, a copper phthalocyanine dye and a disperse dye into an aromatic hydrocarbon solvent by a mixing treatment of the above dyes with a polymer prepared from polypropylene glycol and ethylene oxide having a general formula as shown below:

^{*}Numbers in the margin indicate pagination in the foreign text.

$HO \leftarrow CH_2 CH_2 O \rightarrow_m \leftarrow CHCH_5 CH_2 O \rightarrow_n$

---(CH2CH2O)m1H(1)

(wherein in the above formula, n is integers of 15 \sim 50; m + m' is integers of 5 \sim 30; and the sum of m + m' + n is 20 \sim 80).

Conventionally, as solvent-soluble dyes, 2:1 type metal complex salt dye which is subjected to a salt-forming treatment with an aliphatic amine, an alicyclic amine or a heterocyclic amine (JP-B (Tokko) Sho41-12915 Detailed specification, JP-B (Tokko) Sho44-6597 Detailed specification) or with an aliphatic quaternary ammonium salt (JP-A (Tokkai) Sho48-8316 Detailed specification) have been known. For a copper phthalocyanine /136 dye, that having a sulfonic acid group which is subjected to a salt-forming treatment with an alkylamine [Zabon Fast Blue HFL, from BASF, Senryou Binran (Dye Handbook), new edition, P 865, Edited by the Organic Synthesis Chemistry Society] or with an alkoxyalkylamine (JP-B (Tokko) Sho44-3218 Detailed specification) have been known. For an acid dye, that which is subjected to a salt-formation treatment with an alicyclic amine (Zabon Fast Yellow CGG, from BASF Co., Dye Handbook, P 840, Edited by the Organic Synthetic Chemistry Society) has been known. However, although those dyes are soluble in a polar solvent such as ethanol, acetone, methyl cellosolve, methyl ethyl ketone, ethylene glycol, ethyl acetate, n-butanol and the like, they are almost completely insoluble in an aromatic hydrocarbon solvent such as benzene, toluene, xylene, styrene, cymene, naphthalene and the like. Even if they are soluble, the amount which can be dissolved is very small.

The present inventors have done extensive research concerning the dye which is soluble in an aromatic hydrocarbon solvent and discovered that by a simple physical mixing of the dye with a polymer prepared from polypropylene glycol and ethylene oxide as shown in the general formula (1), the dye exhibits better-than-expected solubility into an aromatic hydrocarbon solvent such as styrene, benzene, toluene, xylene, cymene, naphthalene and the like, which has led to the completion of the present invention.

The present inventors have already submitted patent applications concerning a method of manufacturing a 2:1 type metal complex salt dye which is soluble in a hydrocarbon solvent in JP-A (Tokugan) Sho48-60325 and JP-A (Tokugan) Sho48-44223. Those patent applications have a characteristic in that a sodium salt, a potassium salt, or an ammonium salt of 2:1 type metal complex salt dye is subjected to a salt-formation treatment with a fatty acid salt of N-alkyl-substituted alkylenediamine. Also, in the JP-A (Tokugan) Sho50-28650, the patent application for another solubilization method of 2:1 type metal complex salt dye in the hydrocarbon solvent has been submitted. In this method, an aliphatic ammonium salt or alicyclic ammonium salt of 2:1 type metal complex salt dye is subjected to a mixing treatment with a fatty acid salt of N-alkyl-substituted alkylenediamine. Also, in regard to a method of manufacturing a hydrocarbon solvent-soluble copper phthalocyanine dye, the patent application has been submitted in JP-A (Tokugan) Sho48-59698. In this patent application, a sodium salt, a potassium salt, or an ammonium salt of copper phthalocyaninesulfonic acid is subjected to a salt-formation treatment with a fatty acid salt of N-alkyl-substituted alkylenediamine. Also, the patent application has been submitted concerning another method of manufacturing a hydrocarbon solvent-soluble copper phthalocyanine dye in the JP-A (Tokugan)

Sho50-48653. In this patent application, an aliphatic ammonium salt or alicyclic ammonium salt of copper phthalocyaninesulfonic acid is subjected to a mixing treatment with a fatty acid salt of N-alkyl-substituted alkylenediamine.

In the method of the present invention, dyes can be easily made aromatic hydrocarbon-soluble by a simple mixing treatment of the dye with a polymer prepared from polypropylene glycol and ethylene oxide as shown by the general formula (1).

That is, in the first method of the present invention, a dye can be made easily solubilized in an aromatic hydrocarbon solvent by first dispersing the dye in the aromatic hydrocarbon solvent such as toluene, xylene, cymene, styrene, naphthalene and the like, then, while heating and stirring, adding a polymer prepared from polypropylene glycol and ethylene oxide as shown in the general formula (1) into the dispersion in an amount of at least 100% based on the amount of the dye, or preferably 150 ~ 200%. In this case, by adding the polymer prepared from polyethylene glycol and ethylene oxide, the dye is solubilized in the aromatic hydrocarbon solvent. The obtained solution in which the dye is completely solubilized can be used as it is as a concentrated dye solution or it can be used in a paste form after removing the used solvent.

Furthermore, in the second method, the dye which is soluble in any desired solvent can be obtained easily by simply kneading a mixture of the dye and the polymer prepared from polypropylene glycol and ethylene oxide as shown in the general formula (1) on a 3-roll kneader at room temperature or under heating. In this case, the amount of the polymer prepared from polypropylene glycol and ethylene oxide to be used is at least 100% based on the amount of the dye, or preferably at least 200%. In this case, the dye composition in a paste form which is soluble in the aromatic /13° hydrocarbon solvent can be easily obtained.

As examples of the polymer prepared from polypropylene glycol and ethylene oxide, those having the following structures can be appropriately used:

НО	-{	-сн	2 CH 2 O)m	€C£	1CH	5 CH	₂ O)	п (сн ₂ сн ₂ о) _т , н
1)	b	=	15	то	+	m'	=	5
2)	n	7	20	m	+	m*	452	23
5)	D	~	23	m	+	m'	=	50
4)	71	ex	35	m	+	m '	155	10
5)	n	**	17	m	+	m i	ca	7
6)	n	c#	32	m	+	m ¹	ø	30
7)	n	128	50	m	+	m '	**	18
8)	n	-	27	m	÷	m ¹	=	7
9)	8	=	19	m	+	m,	100	26
10)	n	(CA)	20	យ	+	m '	at	10
11)	n	€	1 4	w	+	m'	ER	9
12)	n	g/3.	18	រា	+	m r	UM.	20 .

As examples of the dye which can be appropriately used in the present invention, 2:1 type chrome and cobalt complex salt dyes [C.I. Solvent Yellow

61 (Aizen Spilon Yellow GRH, from Hodogaya Kagaku Co.), C.I. Solvent Red 83 (Aizen Spilon Red BBH, from Hodogaya Kagaku Co.), C.I. Solvent Black 22 (Aizen Spilon Black BH, from Hodogaya Kagaku Co.) and the like], 1:1 type Chrome and Cobalt complex salt dyes [C.I. Solvent Yellow 19 (Zabon Fast Yellow GR, from BASF Co.) and the like], copper phthalocyanine dye (Zabon Fast Blue HFL, from BASF, and the like), and a disperse dye [C.I. Disperse Red 4 (Diaseriton Fast Pink R, from Mitsubishi Kasei Co.), and the like] can be mentioned.

The form of the aromatic hydrocarbon solvent-soluble dye which can be obtained in the present invention is a dye solution containing the solubilizing agent, a dye paste containing the solubilizing agent prepared by removing the solvent from the above dye solution, or a dye paste prepared by the 3-roll kneader (hereinafter, these will be referred to as the dye composition). The paste form of the dye composition may be, in some cases, mixed with an appropriate solvent to be used as a liquid product having an appropriate concentration depending on the use. This indicates that the method of the present invention can overcome the disadvantages of the powder or flake form of the conventional solvent-soluble dyes, which are the common form of the current solvent-soluble dyes, such as a slow dissolution rate into the solvent, difficulty in handling, only a slight solubility into the aromatic hydrocarbon solvents, extra equipment requirements necessary for dissolution of the dye and the like.

Furthermore, one of the major characteristics of the present invention is the fact that the dye can be made soluble into non-polar aromatic

hydrocarbon solvents by a simple mixing treatment of the dye with the polymer prepared from polypropylene glycol and ethylene oxide. This is very advantageous from the standpoint of industrial application. Also, the fact that the dye composition of the present invention exhibits a larger solubility as compared to the current oil-soluble dye is also one of the major characteristics of the present invention.

The dye compositions obtained by the present invention have wide application areas because of their chemical and physical characteristics. For example, they can be used as a variety of inks and lacquers, as an excellent coloring agent for wood materials, oils, natural and synthetic waxes, synthetic resins and the like, and as a coloring modifier for petroleum products such as a gasoline, a lubricating oil and others.

The present invention will be explained in detail using examples. However, the present invention will not be restricted to these examples. In the examples, the terms "parts" and "%" means "weight parts" and "%" by weight", respectively, unless described otherwise.

Example 1

The 2:1 type chromium complex salt dye having the structure as shown below:

in an amount of 50 parts was dispersed in 200 volume parts of xylene and mixed, while stirring, with 75 parts of the polymer prepared from polypropylene glycol and ethylene oxide having a structure as shown below:

$$HO \leftarrow (CH_2CH_2O)_{\overline{m}} \leftarrow (CH_2CH_2O)_{\overline{m}} + CH_2CH_2O)_{\overline{m}} + H$$

The temperature of the mixture was maintained at 90 ~ 100°C for 2 hours. After confirming that the 2:1 type chromium complex salt dye was completely dissolved, xylene used as the solvent was removed to obtain 130 parts of the dye composition.

This dye composition was soluble in the aromatic hydrocarbon solvent such as toluene, xylene, styrene and the like, showing a yellow color and exhibiting a solubility of 200 parts/100 volume parts of styrene. On the other hand, 2:1 type chromium complex salt dye, prepared without the above treatment, exhibited a solubility of 0.02 parts/100 volume parts of styrene. Example 2

The 2:1 type metal complex salt dye having a structure as shown below:

in an amount of 25 parts was dispersed in 300 volume parts of toluene and mixed with 50 parts of the polymer prepared from polypropylene glycol and ethylene oxide having a structure as shown below:

HO
$$-(CH_2CH_2O)_m$$
 (CH $CH_3CH_2O)_n$ (CH₂CH₂O)_m H

n ≈ 21 , m + m' ≈ 20

The temperature of the mixture was maintained at 75 ~ 85°C for 1 hour. After confirming that the 2:1 type chromium complex salt dye was completely dissolved, toluene as the solvent was removed to obtain 80 parts of the dye composition.

This dye composition was soluble in the aromatic hydrocarbon solvent such as xylene toluene, styrene, and the like, showing a black color and exhibiting a solubility of 250 parts/100 volume parts of benzene. On the other hand, 2:1 type chromium complex salt dye, prepared without the above treatment, exhibited a solubility of 0.01 parts/100 volume parts of benzene.

Example 3

The 2:1 type chromium complex salt dye having a structure as shown below:

in an amount of 60 parts was dispersed in 100 parts of benzene and mixed with 100 parts of the polymer prepared from polypropylene glycol and ethylene oxide having a structure as shown below:

HO
$$-(CH_2CH_2O)\frac{1}{m}$$
 (CHCH₃CH₂O)_m (CH₂CH₂O)_m, H
 $\alpha = 55$, m + m³ ~ 10

The mixture was heated at $50 \sim 60^{\circ}\text{C}$ for 0.5 hours to obtain 260 parts of the dye composition solution.

Example 4

The copper phthalocyanine dye having a structure as shown below:

in an amount of 50 parts was dispersed in 250 parts of benzene and mixed with 80 parts of the polymer prepared from polypropylene glycol and $\frac{139}{139}$ ethylene oxide having a structure as shown below:

HO -
$$(CH_2CH_2O)_{\overline{m}}$$
 - $(CHCH_3CH_2O)_{\overline{m}}$ - $(CH_2CH_2O)_{\overline{m}}$ H

 $n = 17$, $m + m' = 10$

The temperature of the mixture was maintained at 50 ~ 60°C for 1 hour. After confirming that copper phthalocyanine dye was completely dissolved, benzene used as the solvent was removed to obtain 133 parts of the dye composition.

This dye composition was soluble in the aromatic hydrocarbon solvent such as xylene, benzene, toluene and the like, showing a blue color and exhibiting a solubility of 250 parts/100 volume parts of xylene. On the other hand, the above copper phthalocyanine dye, prepared without the above treatment, exhibited a solubility of 0.005 parts/100 volume parts of toluene.

Example 5

The disperse dye having a structure as shown below:

in an amount of 40 parts was dispersed in 400 parts of toluene and mixed with 60 parts of the polymer prepared from polypropylene glycol and ethylene oxide having a structure as shown below:

HO
$$-(CH_2CH_2O)_m$$
 $-(CHCH_3CH_2O)_m$ $+ m' = 10$

The temperature of the mixture was maintained at 80 ~ 90°C for 2 hours. After confirming that the disperse dye was completely dissolved, toluene as the solvent was removed to obtain 100 parts of the dye composition.

This dye composition was soluble in the aromatic hydrocarbon solvents such as xylene, benzene, toluene, styrene and the like, showing a red-orange color and exhibiting a solubility of 270 parts/100 volume parts of xylene. On the other hand, the above disperse dye, prepared without the above treatment, exhibited a solubility of 0.1 parts/100 volume parts of xylene.

The dyes as shown in the tables below were subjected to a mixing treatment with the polymer prepared from polypropylene glycol and ethylene oxide having a structure as shown in the general formula (1) to obtain the aromatic hydrocarbon solvent-soluble dye compositions. The solubility in the tables is the solubility of the dye composition in toluene. The dye composition for the evaluation of the solubility was prepared by removing the solvent from the above dye composition after the completion of the mixing treatment. Also, CuPc means the residue of the copper phthalocyanine group.

1	安悠何	a #2	ポリブコピレングリコールと エテレンオキタイドとの复合句	4 m 5 m	6 kg 6 kg	7 m	8 1 / エン 中の合和	9
	6	CONH N = N - C C-OH; N= N - C N=	$BO \leftarrow CH_{2}CH_{2}O \xrightarrow{m} (CHCH_{2}CH_{2}O)^{m}$, H $B = 31$, $m + m' = 26$	10	90~ 100	1	12 \$##6	3 6 0
	7	C2H6 C2H6 C2H6 C2H6 C2H6 C2H6 C2H6 C2H6	но-{сн ₂ сн ₂ о } (с нсн ₃ сн ₂ о) п (с н ₂ сн ₂ о) п, н п ∞ 25, т 4 m' ≈ 26	11	75~ 80	2	∌ € 13	. 280

- 1. Example
- 2. Dye
- 3. Polymer prepared from polypropylene glycol and ethylene oxide
- 4. Mixing treatment
- 5. Solvent
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Xylene
- 11. Toluene
- 12. Green-yellow color
- 13. Red color

/141

1 %	g g 2	ポリプロピレンクリコールと <mark>③</mark> エテレンオキナイドとの無合物	4 4 5		13 MJ	1/427 400#1	高 帯 政 (数数/トルエン 100 容量係)
8	0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0	но -{CH2 cH2 о} = (онси3 сн3 о <u>3=</u> 	10	50~ 60	3	## [®]	276
9	GPS GHC ⁴ H ³ NHCH ³) ³ CP L C 20 ⁴ H	mo-{ch₂ch₂o}m-{chch₂ch₂o)m _—-(ch₂ch₂o)m, н в 19, m + m' 17	11	79~ 80	1.0	7 6 13	240

Key:

- 1. Example
- 2. Dye
- 3. Polymer prepared from polypropylene glycol and ethylene oxide
- 4. Mixing treatment
- 5. Solvent
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Benzene
- 11. Toluene
- 12. Violet color
- 13. Blue color

ソ

* * 1	2 8	3 ポリプロゼレングリコールと	4 88	章 類 数 2 の 6	13. 19 14 (b) 7	8 77.37 4006	高 別 民 (受称/トルエン 100 を量節)
\$D	C4 C	HC $-(CH_1CH_2O)\frac{1}{m}(CHCH_3CH_2O)\frac{1}{m}$. H $-(CH_3CH_2O)\frac{1}{m}$. H $n = 15$, $m * q' = 14$	10	50~ 60	20	##8 12	190
11	(803 H)3	но-{сн ₂ ся ₂ о дд (спся ₃ си ₂ о) д — {ск ₂ си ₂ о дд в с = 28, сп + сп = 19	11	80~ 90	10	₹ 6 . 13	150

- 1. Example
- 2. Dye
- 3. Polymer prepared from polypropylene glycol and ethylene oxide
- 4. Mixing treatment
- 5. Solvent
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Benzene
- 11. Toluene
- 12. Blue-green color
- 13. Blue color

12	_{#4} 2	ポリプロゼレングラコールと エテレンオキサイドとの重な物 3	4 1	8 K	81 17)	1,3-2.7 BACP 8	路 州 里 (取料/トルエン (取料/トルエン
12	CuPo(sO3H	ко -{С н ₂ сн ₂ о }, (сясн ₃ сн ₂ о), в (ся ₂ ся ₂ о ;, в	10	80~	1.0	12	110
13	CH3 CH3	HO-{CH2CH2O}m-(CHCH3CH2O)m, H	111	6 0~ 6 5	20	85 # E	120

- 1. Example
- 2. Dye
- 3. Polymer prepared from polypropylene glycol and ethylene oxide
- 4. Mixing treatment
- 5. Solvent
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Xylene
- 11. Toluene
- 12. Blue color
- 13. Orange-red color

/144

199	2 *	ポリプロゼレングリコールと エテレンオキサイドとの重要も 3	4 m	+ #5 ** #4	## # 7		お .第 度 (数科/トルエン 100 写彙件)
4	O NH ₃ OCH ₅	но -{сн ₁ ся ₂ о \ 	10	50~ 60	25	∌ € 111	180

Key:

- 1. Example
- 2. Dye
- 3. Polymer from polypropylene glycol and ethylene oxide .
- 4. Mixing treatment
- 5. Solvent
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Benzene
- 11. Red color

Example 15

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The 2:1 type chromium complex salt dye having a structure as shown below:

in an amount of 40 parts was mixed with 80 parts of the polymer prepared from polypropylene glycol and ethylene oxide having a structure as shown below:

HO -
$$(CH_2CH_2O)_m$$
 (CHCH₃CH₂O)_m (CH₂CH₂O)_m, H
n m 51, m + m' = 21

The mixture was kneaded using a 3-roll kneader (from Inoue Manufacturing Co.) at a room temperature for 1 hour to obtain 118 parts of the dye composition.

This dye composition was soluble in the aromatic hydrocarbon solvent such as benzene, toluene, styrene and the like, showing a red color and exhibiting a solubility of 210 parts/100 volume parts of xylene. On the other hand, the above 2:1 type chromium complex salt dye, prepared without the above mixing treatment, exhibited a solubility of 0.001 parts/100 volume parts of xylene.

Example 16

The copper phthalocyanine dye having a structure as shown below:

in an amount of 50 parts was mixed with 75 parts of the polymer prepared from polypropylene glycol and ethylene oxide having a structure as shown below:

HO
$$+$$
 CH₂ CH₂ O)_m (CHCH₃ CH₂ O)_m (CH₂ CH₂ O)_m, H

 $= 18$, $m + m^2 = 5$

The mixture was kneaded using a 3-roll kneader at a room temperature for 1 hour to obtain 123 parts of the dye composition.

This dye composition was soluble in the aromatic hydrocarbon solvent such as benzene, toluene, xylene, styrene and the like, showing a blue color and exhibiting a solubility of 250 parts/100 volume parts of toluene. On the other hand, the above copper phthalocyanine dye, prepared without the above mixing treatment, exhibited a solubility of 0.003 parts/100 volume parts of toluene.

Example 18

The disperse dye having a structure as shown below:

in an amount of 20 parts was mixed with 30 parts of the polymer prepared from polypropylene glycol and ethylene oxide having a structure as shown below:

HO
$$-(CH_2CH_2O)_{\overline{m}}$$
 $+(CH_3CH_2O)_{\overline{m}}$ $+(CH_2CH_2O)_{\overline{m}}$ $+(CH_2CH_2O)_{\overline{m$

The mixture was kneaded using a 3-roll kneader at room temperature for 1 hour to obtain 48 parts of the dye composition.

This dye composition was soluble in the aromatic hydrocarbon solvent such as benzene, toluene, xylene, styrene and the like, showing a red color and exhibiting a solubility of 280 parts/100 volume parts of toluene. On the other hand, the above copper dispersed dye, prepared without the mixing treatment, exhibited a solubility of 0.2 parts/100 volume parts of toluene.

The dyes as shown in the following tables were mixed with the polymer prepared from polypropylene glycol and ethylene oxide having a /146 structure as shown by the general formula (1) and the mixture was subjected to a mixing treatment using a 3-roll kneader to obtain the dye compositions which were soluble in the aromatic hydrocarbon solvent. The results are shown in tables below. In the tables, the solubility of the obtained dye compositions was measured using toluene. Also CuPc means the residue of the copper phthalocyanine group.

15	2	3	4 リプロセレンタリコールと エチレンオキサイアとの配合物	3	5 s + 2 g g g	#0 #1 10 13 10 17	8 1227 7060	の 新 変 (東京都立法 100万里式han
19	OgN N = N - NHg OgN NH Og NAG	80	но -{ch ₂ ch ₂ o) _m -{chc ₃ ch ₂ o) _m {ch ₂ ch ₂ o) _m , н а = 17, с • m' = 7	\$ 0	20~ 25	2	80 th 10	250
20	O OH OCH,	40	КО-{CH ₂ CH ₂ O} _m -{CH ₂ H ₂ CH ₂ O) _m ; К (CH ₂ CH ₂ O) _m ; К л ≈ 26, м + м' ≈ 19	45	50~ 55	1	# 6	175

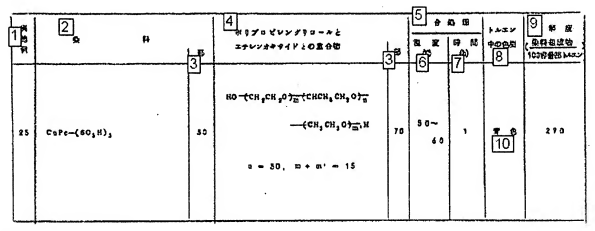
- 1. Example
- 2. Dye
- 3. Parts
- 4. Polymer prepared from polypropylene glycol and ethylene oxide
- 5. Mixing treatment
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Brown color
- 11. Red color

1 %	2	31_	4 リブロビレンタリコールと まナレンオキサイグとの重や物	3	5 a s s s s s s s s s s s s s s s s s s	22 12 R 10 (9) 7	8 }**** *08 %	多年的成立
21	N=N-C ⁶ H ⁴	35	но (сн ₂ сн ₂ о) 	70	Los	. 1	# 6	520
2.2	CH3 CH3	50	р = 30, ф + ш, р 11 (сн³сн³о) ^ш .п но(сн³сн³о) ^ш (сиса³сн³о) п	110	20~ 25	as	2 6	298

- 1. Example
- 2. Dye
- 3. Parts
- 4. Polymer prepared from polypropylene glycol and ethylene oxide
- 5. Mixing treatment
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Brown color

	2 8		4 オフロゼレンタリコーンと エテレンまやすくどとの重点回	3	5 p	107 100 177	+264 +264 8	9 85 R 20 F R C to 100 F 20 1000
	The state of the s	70	HO-(CH ₃ CH ₂ O) m (CHCH ₃ CH ₂ O) m , H	4 50	40~ 45	7	3 2	25 ŋ `
04	C ₂ H ₃ 30 ₂ N C ₂ H ₃ C ₂ H ₃ (30 ₃ H) ₂		ко —{ он, он, от — сыси, си, от п —— (сы, си, от п, и п = 52, м + м, м 23	SO	20~	1.5	* & &	223

- 1. Example
- 2. Dye
- 3. Parts
- 4. Polymer prepared from polypropylene glycol and ethylene oxide
- 5. Mixing treatment
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Yellow color
- 11. Blue color



- 1. Example
- 2. Dye
- 3. Parts
- 4. Polymer prepared from polypropylene glycol and ethylene oxide
- 5. Mixing treatment
- 6. Temperature (°C)
- 7. Time (h)
- 8. Color in toluene
- 9. Solubility (dye/100 volume parts toluene)
- 10. Blue color